

DEPARTMENT OF MICROBIOLOGY
EUCLID AVENUE AND KINGSHIGHWAY

Dear La:

the proofs arrived last week and have been not returned. The reprint order is enclosed.

I have been able to get a little work done since we stopped teaching (which turned out to be a much greater (se more tring) chare than anticipated). If currently, I am trying to finish up the UV effect on recombination. I got some closely linked t's (T4) from Benzer and four found that starting at a control recombination value of 10-3 the value varied himsely with the UV done as for as I went (n4 fold increase, P/Po~10-5). Hering this pair of t's, the UV increases the frequency (also the absolute number) of cells which liberate any recombinants at all. I take this as good evidence that UV does not

merely cause a selection for recombinants, produced at the no control rate, but rather increases the number of recombinations. I will talk these things over with Seymour when he visits this week. In any event, all I will do along these lines is a ten more crosses of this type, and some crosses to see the effect of UV on the number of rounds of mating. I will write a short report of the fundings and stop, since I don't see a further development of the work.

Much of these results and those on crossreactivation would be easier to think about if
we had information on the multiplication of the
dead phages perse. I am setting up to see
if cells singly infected with UVT2 and P32
pilled T2 can make DNA with HMC. in
we will look for the small amounts
of HMC expected by infecting in a medium
containing C'Y (abelled queene or acetate and
looking for C'Y in the HMC spot on chromatograms. Probably this will turn out to be mainly
a biochemical exercise for me (eq. making
C'Y queene) but that is all right, too.

SCHOOL OF MEDICINE SAINT LOUIS

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A further example of my diddlings is my attempt last week to get naked phage DMA into cells. Since things!

I think it will work with a little trick to be tried tomorrow. I will let you know if anything happens and only mention it here because at (ast someone has tried it. The idea is to try and find out something about the physical nature of the three linkage groups.

I will try to come to Unbana the third week in April on my way East and will book torward to seeing you then. Please remember me to Zella and to everyone in the (ab. we were all sorry to hear of Sol's accident and qlad to learn he was back so soon.

I think I will describe the phage DHA experiment in the hope that you will have some suggestions to make. The DHA is prepared by osnotic shock from T2h Pf (Pf = proflowing resistant). And is carefully cleaned by centralyation

and antiphoge serum so that the effect of ghosts are absent and the background of line phages is absolute zero. The preparation I have has 10 mpphage equivalents of DMA and almost resembles egg white in physical properties. I expose B to it at 37° after interting with T2 F13 at 0° and high multiplicity, to maximize the usefulness of the leakiness who of the cells. After serum treatment I plate the cells on S(x). 13 interted cells do not plate but if the DNA got in (it is t+) they might. The check would be in testing any plagnes for the other markers in the multiply marked MA. The experiment is spouled by the high background of mutants in all my TIT Stocks which can plate on s(a). UVing them does not help much because of MR. I am X- raying tomarion. My TYT stocks have the neckessary low background and I will also try there with the TZ DNA. A success m this experiment would be pretty and I would appreciate any thing you have to offer on the idea. with best regards -

Bob